Molecular and Cellular Differences in Cardiac Repair of Male and Female Mice

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ORIGINAL RESEARCH

Molecular and Cellular Differences in Cardiac Repair of Male and Female Mice

Amanda B. Pullen, BS; Vasundhara Kain, PhD; Charles N. Serhan, PhD, DSc; Ganesh V. Halade, PhD

BACKGROUND: Leukocyte-directed biosynthesis of specialized proresolving mediators (SPMs) orchestrates physiological inflammation after myocardial infarction. Deficiency of SPMs drives pathological and nonresolving inflammation, leading to heart failure (HF). Differences in SPMs and inflammatory responses caused by sex-specific differences are of interest. We differentiated leukocyte-directed biosynthesis of lipid mediators in male and female mice, focusing on leukocyte populations, structural remodeling, functional recovery, and survival rates.

METHODS AND RESULTS: Risk-free male and female C57BL/6 mice were selected as naïve controls or subjected to myocardial infarction surgery. Molecular and cellular mechanisms that differentiate survival, heart function, and structure and leukocyte-directed lipid mediators were quantified to describe physiological inflammation after myocardial infarction. Female mice show improved survival in acute HF but no statistical difference during chronic HF compared with male mice. Female mice improved survival is marked with functional recovery and limited remodeling compared with male mice. Male and female mice are similarly responsive to arachidonate lipoygenase (LOX-5, LOX-12, LOX-15) or cyclooxygenase (COX-1, COX-2) in acute HF and particularly male infarcted heart had overall increased SPMs. Female cardiac healing is marked with the biosynthesis of differential p450-derived product, particularly 11,12 epoxyeicosatrienoic acid in acute HF. A sex-specific difference of dendritic cells in acute HF is distinct, with limited changes in chronic HF.

CONCLUSIONS: Cardiac repair is marked with increased SPM biosynthesis in male mice and amplified epoxyeicosatrienoic acid in female mice. Female mice showed improved survival, functional recovery, and limited remodeling, which are signs of fine-tuned physiological inflammation after myocardial infarction. These results rationalize the sex-specific precise therapies and differential treatments in acute and chronic HF.

Key Words: heart failure ■ ischemia ■ leukocytes ■ resolution of inflammation ■ specialized proresolving mediators

Cardiovascular disease (CVD) is a leading cause of death in the United States, with close to one third of deaths being attributed to some form of cardiometabolic pathological feature.1,2 The primary risk factors for CVD are related to several lifestyle aspects that increase the likelihood of myocardial infarction (MI; heart attack) in men and women. MI events are less common in premenopausal women and occur in men at an earlier age (≈65 years) compared with women (≈72 years).3-5 The epidemiological study on the autonomic response to CVD compounded with obesity showed, although men and women are both negatively impacted by obesity, some cellular and molecular mechanisms of these responses are sex specific and the overall mechanism is incomplete.6 There are differences in the risk, timing, and symptoms of heart disease between men and women for a variety of pathological conditions; however, estrogen has been linked to a higher survival rate of cardiomyocytes after MI.7 Sex-specific differences in cardiovascular health stem from multiple factors, including leukocytes, fat immune metabolism, glycoprotein content, gene expression, sex hormones, and sociocultural aspects, like behavior, environment, and nutrition.8-11
In response to cardiac injuries, such as MI, the leukocyte-directed initiation of inflammation and activation of resolution programs facilitates the clearance and repair of damage myocardium. Delayed initiation of inflammation or immune suppression causes the late arrival of leukocytes to the site of infarction and prolonged residual time of leukocytes during the clearance process; when polymorphonuclear neutrophils are not cleared on time, it can lead to increased, chronic inflammation. After experimentally inducing MI by coronary ligation, leukocytes biosynthesize endogenous specialized proresolving mediators (SPMs) that are essential for cardiac healing and repair. SPMs are lipoxygenase-mediated omega 3 fatty acid derived biomolecules, including maresins, protectins, resolvins, and lipoxins, that have been shown to have a feed-forward effect on the adaptive immune system, leading to potential therapeutic uses to control inflammation and alleviate chronic diseases. A positive-feedback loop between SPMs and leukocytes encourages the resolution of injury-induced inflammation by increasing differential lipoxygenase activity. Differences in SPMs between men and women are still being uncovered; therefore, we propose to investigate the differences of cardiac healing in male and female mice with major emphasis on leukocyte trafficking, endogenous bioactive lipid mediators (including SPMs), heart structure, function, and post-MI survival in acute heart failure (AHF) and chronic heart failure (CHF). In the presented report, we aim to explain the differences of male and female C57BL/6J mice in 3 key aspects: (1) the signaling of lipoxygenases and/or cyclooxygenases; (2) the biosynthesis of SPMs in regard to cardiac healing and repair; and (3) cardiac healing in the context of physiological inflammation and resolution response. Our report suggests that male and female mice operate cardiac repair with the biosynthesis of SPMs differently. Male mice have increased overall SPM levels, and female mice have amplified epoxyeicosatrienoic acids (EETs) in AHF with increased levels of reparative macrophages.

**METHODS**

Data presented in this article support the findings of this study and are available from the corresponding author on reasonable request.

**Animal Compliance**

All animal procedures were conducted according to the National Institutes of Health Guide for the Care and Use of Laboratory Animals (eighth edition, 2011) and were approved by the Institutional Animal Care and Use Committees of University of South...
High-Resolution Transthoracic Echocardiography

In vivo cardiac performance of mice was assessed using high-resolution noninvasive echocardiography (Vevo 3100; VisualSonics, Canada) during which mice were anesthetized using a 1-2% isoflurane and 100% oxygen mixture, administered with a small rodent respirator. Body hair on the dorsal side of each animal was removed using depilatory cream at least an hour before the imaging (Nair). Echocardiographic measurement of cardiac structure and function was performed with the high-resolution echocardiography analysis system Vevo 3100 equipped with MX 400 transducer (30 MHz) for small animals. Each measurement was done by a single trained technician (A.B.P.) to eliminate interpersonal variation. Two-dimensional B-mode short-axis and long-axis views, along with B-mode tracings of the left ventricle (LV), were obtained. Using analysis software provided by the manufacturer, the following data were obtained: LV dimension at systole and diastole, posterior wall thickness at systole and diastole, interventricular septal wall thickness at systole and diastole, LV volume at systole and diastole, LV stroke volume, ejection fraction, fractional shortening, heart rate, cardiac output, global longitudinal and global circumferential strain, and LV mass. In addition, the integrated monitoring system allowed measurement of physiological parameters, such as body temperature, ECG, heart rate, and respiration, to be captured and monitored throughout an imaging session.20

Coronary Ligation Microsurgery to Induce MI and Irreversible HF

The hair on the dorsal side of the animal was removed before the surgery. Mice were anesthetized with a 2% isoflurane and 100% oxygen mixture and secured to the operating table. The MI was induced in the mice via ligation of the left anterior descending artery using minimally invasive surgery, as previously described.21 A total number of 71 mice were subjected to this surgery.20

Measurement of Post-MI Survival

After coronary ligation, mice were monitored closely for recovery and survival monitor and classified into 3 categories. Perioperative mortalities included animals that died during surgery or within the first 24 hours after surgery. Mice that survived 24 hours after surgery but died within 7 days were included within the AHF survival curve. Mice that survived at least 8 days after surgery but died before day (d) 56 were considered in the CHF survival curve. A necropsy was performed on the mice that died after 24 hours to determine if the mortality was caused by a rupture of the LV or by CHF. Ruptures included all mice with a tear or hole on the LV, resulting in clotted blood within the chest cavity. CHF deaths included mice with no apparent tear or clotted blood in the chest cavity.4,20,22

Necropsy

Animals were anesthetized using a 2% isoflurane and 100% oxygen mixture. From each animal, the LV, lungs, spleen, and tibia were collected. The harvested tissues were either flash frozen and stored at −70°C and used for Reverse transcription polymerase chain reaction and/or Western blot analysis or placed in 10% zinc formalin and transferred to 70% ethanol after 24 hours and used for histological analysis. The right ventricle was removed from the LV and weighed. The LV was then separated into 3 pieces: the infarct area (infarcted LV [LVI]) or ischemic area below the ligation, the middle piece or the area that has both infarct and remote areas, and the remote area (remote LV) or the area not yet severely affected by the ligation. The ischemic area of the LV and a portion of the spleen were used for gene expression and histological analysis and compared with naïve controls. The lungs are weighed and then placed in an incubator (37°C) for 24 hours, and dry weight is recorded.20

Histological Analysis: Hematoxylin and Eosin Staining

Sections of the LV at No-myocardial infarction (MI), MI- d1, and MI- d56 were stained using hematoxylin and eosin, and images were acquired using a BX43 microscope with an attached Olympus DP73 camera, as previously described.20

Confocal Microscopy for Fibrotic Remodeling

For immunofluorescence, sections of the LV at MI- d5 were stained using α-smooth muscle actin, discoidin domain receptor 2, and Hoechst; and confocal images were acquired using a Nikon A1 high-resolution microscope, as described previously.23

Leukocyte Quantitation Using Flow Cytometry

The LV and spleen of No-MI (d0), MI- d1, MI- d3, and MI- d56 male and female mice were taken; and mononuclear cells from both were isolated using methods as previously described.24 After adjusting for ≈1 to 2 million cells/stain, the cells were suspended in...
100 μL of FC block and allowed to incubate for 10 minutes on ice. A stain cocktail containing fluorophore-labeled monoclonal antibodies in 2× concentration was added on ice for 30 minutes. The stain cocktail contained CD45-PE-CY7 (BD Biosciences, San Jose, CA), CD11b-APC, F4/80-PERCP (Molecular Probes, Eugene, OR), Ly6C-FITC (BD Biosciences), Ly6G-Pacific blue (e-Bioscience), and LIVE/DEAD Fixable Blue Dead Cell Stain Kit and was used to determine viability and cell type (Figure S1A; gating strategy).12,25

Quantitative Real-Time Polymerase Chain Reaction of Gene Transcripts
RNA was isolated from both the LV and spleen of day 0 (d0) and MI-day 1 (MI-d1) male and female mice. Reverse transcription was performed with 2.0 μg of RNA using the SuperScript VILO cDNA Synthesis Kit (Invitrogen, CA). The gene expression levels of COX-1, COX-2, ALOX-5, ALOX-12, ALOX-15, IL-1β, TNF-α, Arg-1, Mrc-1, ALX/FPR2, EP2, EP4, and EPHX-2 were measured using TagMan Probes (Applied Biosystems, CA). The expressions were normalized using hypoxanthine phosphoribosyltransferase 1 (HPRT1). The results were reported using 2−∆Ct values.20

Protein Extraction and Immunoblotting
Protein was extracted from the LVI or the remote LV, as previously described.26 After electrophoresis of 5 to 10 μg of LVI or remote LV protein, primary antibodies ALOX-5, HO-1, COX-2, ALX/FPR2, TNF-α, or GAPDH were allowed to incubate overnight at 4°C. After the incubation, respective secondary antibody (1:10 000 dilution in a 5% blocking solution) was added, as previously described.26 Femto chemiluminescence detection system was used to detect protein expressions (Pierce Chemical, Rockford, IL). To normalize the total protein/lane, densitometry was performed using Image J software (NIH, Bethesda, MD).26

Quantification of SPMs Using Mass Spectrometry
Samples of the LV and spleen for liquid chromatography–tandem mass spectrometry analysis were obtained using solid phase extraction columns, as described.72 To observe and measure the levels of lipid mediators in the LVI, an LC-tandem MS/MS system, QTrap 5500 (ABSciex), equipped with an Agilent HP1100 binary pump, was used. The Agilent Eclipse Plus C18 column (50 mm×4.6 mm×1.8 μm or 100 mm×4.6 mm×1.8 μm) with a gradient of methanol/double-distilled H2O/acetic acid (60:40:0.001 to 100:0:0.01) was used at a flow rate of 0.5 mL/min. If a standard for a product was not available, a product with similar chromatographic behaviors was used for a calibration curve. Deuterium-labeled internal standards included lipoxin A4, resolvins D1 (RvD1), Prostaglandin E2 (PGE2), leukotriene B4, and others, obtained from Cayman Chemical Company (Ann Arbor, MI) and used to calculate recoveries. The peak area of each Multiple Reaction Monitoring (MRM) transition and linear calibration curves were used for the quantification of each mediator.27 A minimum of 6 characteristics and diagnostic ions were used for identification of all SPMs, in accordance with published criteria of biologic and synthetic mediators.27,28

Statistical Analysis
The results are expressed as mean±SEM. Analyses were performed using a 2-way ANOVA (mixed model); multiple comparisons are done using Sidak test recommended parameter for time kinetics and post Tukey test for other comparisons (Graphpad Prism 8.3.0 application). The Kaplan-Meier test and log-rank test were followed for survival analysis. P<0.05 was considered as statistically significant.

RESULTS
Female Mice Have Higher Survival With Few Ruptures in AHF
To determine the sex-specific differences between male and female mice after permanent coronary ligation, we measured long-term survival, cardiac function, myocardium structural integrity, and gene and protein expression (Figure 1A; study design). A total of 43 male mice and 28 female mice were subjected to occlusion of the left anterior descending coronary artery. Successful MI and LV dilation were confirmed using echocardiography within 24 hours after surgery with the inclusion criteria of fractional shortening <15%. Any mortalities within the first 24 hours were considered perioperative mortalities and excluded from the survival curve (=5% for the trained surgeon). After analyzing the mortality rate of the mice after MI using Kaplan-Meier curve, we separated the deaths into either AHF (MI-d1 to MI-d7) or CHF (MI-d8 to MI-d56). Deaths in these categories were further divided into HF or ruptures. In response to coronary ligation, the overall survival in female mice is 21% higher compared with male mice, without developing CHF or a rupture up to 56 days after MI (Figure 1B and 1C). A breakdown of the survival chart into the short- and long-term phases shows that male mice have increased mortality attributable to ruptures (28% compared with 7% in female mice) and CHF (12% compared with 7% in female mice).
Within 7 days of MI (Figure 1D; Figure S1B). Within the long-term phase (after MI-d7), there is no significant difference in either rupture or CHF mortalities between male and female mice (Figure 1E; Figure S1C). This in-depth survival analysis allowed for the more accurate differentiation between male and female survival after MI and pinpointed the sex-specific responses to AHF and CHF.

Female Mice Show Improved Recovery and Heart Function After MI
Gravimetric parameters were measured to more accurately rule out variables between male and female mice (Table 1). After the coronary ligation surgery, echocardiograms of both male and female mice showed reduced cardiac function consistent with HF (Figure 2). In both sexes, LV dilation was evident by the decrease in vector length (longitudinal and circumferential axis b-modes), increased strain on the myocardium (strain), and myocardium dysynchronicity (segmental synchronicity) of the representatives for MI-d1 to MI-d56 compared with the No-MI controls (Figure 2). However, after an initial decrease, female mice recovered a portion of their cardiac function at MI-d3, which continued to gradually increase until MI-d56, shown by increased vector length (longitudinal and circumferential axis b-modes), decreased muscle strain (strength), and improved synchronicity (segmental synchronicity) (Figure 2). The improvement of the female mice heart function is further shown by the increase in their fractional shortening (MI-d56: 19% in females compared with 8% in males) and global longitudinal strain (MI-d56: −11% in females compared with −6% in males) (Table). Thus, improvement in heart functional recovery is believed to be enabling the female mice to “bounce back” and survive at a significantly higher rate than male mice after MI.

Cardiac Tissue Repair Marked With SPM Biosynthesis With Higher Levels of EETs in Female Mice
After discovering the higher survival in females is caused by lower rupture rates in AHF, we quantified the levels of proinflammatory mediators, endogenous SPMs, and EETs in the LVI at d1 after MI (Figure 3A through 3F and Figure S2). The arachidonic acid–derived proinflammatory mediators, such as leukotriene B4, Prostaglandin (PG)D2, PGE2, PGF2α, and Thromboxane B2 (TXB2), were increased in both male and female mice as signs of initiation of an inflammatory response (Figure S3) and simultaneous biosynthesis of SPMs as resolving response (Figure S4).
Table. Echocardiography and Necropsy Parameters for Male and Female Mice at MI-d1, MI-d3, MI-d5, and MI-d56 Compared With Naïve Controls.

<table>
<thead>
<tr>
<th>Echocardiographic Parameters</th>
<th>No-MI Controls</th>
<th>MI-d1</th>
<th>MI-d3</th>
<th>MI-d5</th>
<th>MI-d56</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male (n=18)</td>
<td>Female (n=18)</td>
<td>Male (n=19)</td>
<td>Female (n=19)</td>
<td>Male (n=5)</td>
</tr>
<tr>
<td>Heart rate, bpm</td>
<td>48±8±7</td>
<td>475±6</td>
<td>525±17</td>
<td>500±13</td>
<td>568±27</td>
</tr>
<tr>
<td>EDD, mm</td>
<td>4±0.09</td>
<td>4±0.08</td>
<td>4±0.09</td>
<td>4±0.07</td>
<td>5±0.19</td>
</tr>
<tr>
<td>ESD, mm</td>
<td>3±0.11</td>
<td>2±0.08</td>
<td>4±0.12*</td>
<td>3±0.11*</td>
<td>4±0.18*</td>
</tr>
<tr>
<td>Fractional shortening, %</td>
<td>36±2</td>
<td>35±1</td>
<td>10±1*</td>
<td>13±2*</td>
<td>13±3*</td>
</tr>
<tr>
<td>IVSd, mm</td>
<td>0.82±0.04</td>
<td>0.93±0.05</td>
<td>0.71±0.07*</td>
<td>0.70±0.05*</td>
<td>0.62±0.08*</td>
</tr>
<tr>
<td>PWTD, mm</td>
<td>0.75±0.02</td>
<td>0.78±0.04</td>
<td>0.64±0.04*</td>
<td>0.66±0.04*</td>
<td>0.68±0.07*</td>
</tr>
<tr>
<td>IVSs, mm</td>
<td>1.16±0.05</td>
<td>1.16±0.07</td>
<td>0.82±0.08*</td>
<td>0.78±0.05†</td>
<td>0.81±0.05*</td>
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<tr>
<td>PWTs, mm</td>
<td>1.06±0.06</td>
<td>0.99±0.03</td>
<td>0.74±0.05*</td>
<td>0.80±0.06†</td>
<td>0.80±0.05*</td>
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<tr>
<td>GLS</td>
<td>−20±1</td>
<td>−20±1</td>
<td>−7±1*</td>
<td>−8±1*</td>
<td>−10±1*</td>
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</tbody>
</table>

Necropsy parameters

<table>
<thead>
<tr>
<th></th>
<th>No-MI Controls</th>
<th>MI-d1</th>
<th>MI-d3</th>
<th>MI-d5</th>
<th>MI-d56</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male (n=18)</td>
<td>Female (n=18)</td>
<td>Male (n=19)</td>
<td>Female (n=19)</td>
<td>Male (n=5)</td>
</tr>
<tr>
<td>Body weight, g</td>
<td>26±1</td>
<td>22±1</td>
<td>26±1</td>
<td>22±1</td>
<td>27±1</td>
</tr>
<tr>
<td>LV, mg</td>
<td>98±7</td>
<td>80±2</td>
<td>100±2*</td>
<td>74±2*</td>
<td>121±8*</td>
</tr>
<tr>
<td>LV/BW, mg/g</td>
<td>4±0.2</td>
<td>4±0.1</td>
<td>4±0.1</td>
<td>3±0.1</td>
<td>4±0.2</td>
</tr>
<tr>
<td>RV, mg</td>
<td>18±0.7</td>
<td>14±0.6</td>
<td>16±0.4</td>
<td>15±1</td>
<td>21±2</td>
</tr>
<tr>
<td>Spleen, mg</td>
<td>89±3</td>
<td>86±2</td>
<td>70±2</td>
<td>72±3</td>
<td>106±5*</td>
</tr>
<tr>
<td>RV mass/BW, mg/mg</td>
<td>0.7±0.03</td>
<td>0.6±0.03</td>
<td>0.6±0.02</td>
<td>0.7±0.03</td>
<td>0.8±0.05</td>
</tr>
<tr>
<td>Lung mass/BW, mg/mg</td>
<td>7±0.6</td>
<td>7±0.5</td>
<td>7±0.3</td>
<td>7±0.3</td>
<td>7±0.4</td>
</tr>
<tr>
<td>Tibia, mm</td>
<td>15±0.2</td>
<td>15±0.2</td>
<td>15±0.2</td>
<td>15±0.2</td>
<td>17±2*</td>
</tr>
</tbody>
</table>

Data are given as mean±SEM. Bpm indicates beats per minute; BW, body weight; EDD, end diastolic dimension; ESD, end systolic dimension; GLS, global longitudinal strain; IVSd, intraventricular septum diastole; IVSs, intraventricular septum systole; LV, left ventricle; PWTD, posterior wall thickness diastole; PWTs, posterior wall thickness systole; RV, right ventricle; MI, myocardial infarction; and day, d. *P<0.05 vs No-MI naïve controls. †P<0.05 vs male mice at respective time point.
pie chart distribution of SPMs reflects high SPM cluster in male mice (17%) compared with female mice (11%) in LVI (Figure 3C and 3D). The individual quantification of D-series resolvins, maresins, protectins, and arachidonic acid–derived SPMs suggests male and female mice biosynthesis of SPMs with activation of proinflammatory mediators, suggestive of overlapping inflammation-resolution response in cardiac repair (Figure 3A through 3E). The female mice have higher levels of p450–mediated arachidonic acid–derived EET compared with the male counterparts, specifically a significant (1.3 fold; \( P < 0.05 \)) increase in 11,12-EETs (Figure 3F). The quantitative analyses of endogenous bioactive lipids suggest that despite similar SPM contributions in cardiac repair, females have the tendency to produce higher levels of arachidonic acid–derived cypoxins (EETs) compared with the males in infarcted LV in AHF after MI.

**Differential Lipoxygenase and Cyclooxygenase Activation in Male and Female Mice in AHF**

After myocardium injury, cyclooxygenase (COX-1 and COX-2) and arachidonate lipoxygenase (ALOX-5, ALOX-12, and ALOX-15) activity increases at the site of infarction. To determine the differences in initiation of inflammation-resolution between male and female mice, the expression of cyclooxygenases and lipoxygenases was quantified from the spleen and LV for the following time points: No-MI, MI-d1, and MI-d56 (Figure 4A through 4M). At MI-d1, ALOX-5 had higher expression in the male spleen compared with females (Figure 4A). In response to cardiac injury, immune responsive lipoxygenases (5, 12, and 15) and cyclooxygenases (1 and 2) are amplified at MI-d1 in both the LV and spleen as signs of inflammatory and beginning of resolution signal for the biosynthesis of SPMs (Figure 4A through 4J). Female splenic and LVI expressions of COX-1 are higher than male expression at MI-d1 (Figure 4G and 4I). The expression levels of lipoxygenases and cyclooxygenases in the spleen and LV of male and female mice returned to the normal, homeostatic levels at MI-d56 as a signal of physiological control and resolved inflammation (Figure 4A through 4J). LV protein levels show male mice produce more ALOX-5 and COX-2 at MI-d1 compared with female mice (Figure 4K through 4M). These results indicate the spleen and...
infarcted heart coordinate endogenous biosynthesis of lipid mediators in cardiac repair during acute myocardial injury.

Distinct Reparative Structural Remodeling in Female Mice in AHF

Adverse myocardium structural remodeling is a primary risk factor for CHF after MI. To define recovery, we examined hematoxylin and eosin stained LV sections at No-MI controls, MI-d1, and MI-d56 (Figure 5A). For each MI time point, we examined the remote (noninfarcted area after MI), peri-infarct (the area between the remote and infarct), and the infarct (thin-wall area of the myocardium) areas to get an accurate representation of the progression of cell apoptosis and cardiac repair. The remote areas at the No-MI and MI-d1 time points of both the male and female mice show parallel cardiomyocyte, elongate, and centrally located nuclei and intercalated disks in myocardium structure (Figure 5A). However, at MI-d56, the tissue damage has expanded to the remote area and the peri-infarct and infarct areas, with persistent scar within the myocardium structure, and the non-myocyte nuclei are misplaced and are congested within the dead tissue, decreasing structural integrity and aligning with the CHF phenotype (Figure 5A). No difference was observed in the cardiomyocyte area between male and female mice (Figure S5). Fibrotic remodeling further confirmed by α-smooth muscle actin and discoidin domain receptor 2 staining to determine fibroblast activation and presence of collagen in AHF at MI-d5 (Figure 5B). The female LVI showed lower expression of α-smooth muscle actin and discoidin domain receptor 2 compared with male LVI, indicating limited fibrotic remodeling (Figure 5B). Thus, limited fibrotic and structural remodeling in female mice supports the improved functional recovery and survival in both AHF and CHF.

Reparative Monocytes Are Higher in Female Mice Compared With Male Mice After MI

Initiation of inflammation in AHF is defined by the entry of leukocytes at the site of injury with continuum phenotype spectrum; therefore, we analyzed how sex-specific differences impacted LV leukocyte populations after MI using flow cytometry during AHF and CHF<sup>24</sup> (Figure S6). At naïve homeostatic setting, male
and female mice show no difference in LV monocyte (CD45+/CD11b+) populations. In response to ischemic insult, male and female monocytes consistently amplified from d1 to d5 after MI. In both male and female mice, monocyte populations declined at d56 (CHF), comparative to pre-MI naïve state. Female mice showed lower monocytes (53±4.3%; P<0.05) when compared with the male mice (68±3.9%) in AHF (after MI-d1, Figure 6A through 6G). Although, in both male and female mice, monocyte population declined at d56 after MI (CHF), in females, monocyte population was significantly higher (16.0±2.1%) compared with the male counterpart, which displayed (1.6±0.2%) of monocytes (Figure 6A and 6E). However, no sex-specific changes in CD11b+ expression were observed (Figure 6B). Furthermore, the phenotypic analysis of LV monocytes on basis of Ly6Clo and Ly6Chi displayed female mice had a higher Ly6Clo population (18±0.7%), compared with male mice (14±0.7%) at d1 after MI, which remained significantly higher in CHF (Figure 6C, 6D, 6F, and 6G). Increased reparative monocytes during AHF support the improved cardiac repair, functional recovery, and limited fibrotic remodeling of female mice.

Female Mice Have Intensified Dendritic Cell Activation in Cardiac Repair Than Male Mice After MI

After the ischemic insult, dendritic cells (DCs) are an essential part of the innate immune leukocyte system. DCs act as a bridge to the adaptive immune system and play a tissue reparative role in cardiac healing, as the inadequate levels of DCs led to increased rupture and impaired reparative fibrosis after MI.29 In the absence of cardiac injury (No-MI), no difference was observed in CD11c-positive cells between male and female mice (Figure 7A and 7E). In response to coronary ligation, there was a consistent increase in CD11c cells in AHF (MI-d1, MI-d3, MI-d5) and a CD11c+ cell population decline at CHF (MI-d56), irrespective of sex-specific. At post-MI-d1, female mice displayed a higher percentage of CD11c+ cells (3.2±0.5%) compared with male mice (1.8±0.2%), and the DC population peaked at d5 and remained significantly higher in females. Compared with AHF, the DC population declined with time in CHF but relatively remained higher in females compared with males (Figure 7A, 7B, and 7C).
7E). As expected, the neutrophils are the first phagocytic responder in cardiac injury. The quantitative analyses of neutrophil populations from male and female mice showed that neutrophils peaked at post–MI d3 and reached the homeostatic level at d56 after MI. The percentage neutrophil population showed no statistical difference in male and female mice in the infarcted area (Figure 7C, 7D, and 7F).

Analysis of macrophage kinetics (CD45<sup>+</sup>CD11b<sup>+</sup>F4/80<sup>+</sup>) showed a peak at MI d5, with a decline in their population at MI d56. However, male and female mice displayed a similar percentage of macrophage populations (Figure S6A through S6E). The macrophages are subclassified as Ly6C<sup>lo</sup> and Ly6C<sup>hi</sup>, indicative of their proinflammatory and/or proreparative function, so we quantified F4/80<sup>+</sup> cardiac macrophages after MI by Ly6Clo/hi expression. Males and females equally amplified the Ly6C<sup>hi</sup> population, but females displayed a higher population of Ly6C<sup>lo</sup> cells (8.5±1.3%) compared with males (1.1±0.1%) at post–MI d56, contributing to improved cardiac repair (Figure S6B and S6F). Leukocyte-directed physiological inflammation in the female is marked with activation of reparative monocytes and macrophages, with an amplified density of DCs in short-term cardiac repair with the limited difference in CHF. This male and female leukocyte diversity suggests physiological cardiac repair in females expedited by the presence of a higher number of DCs and reparative monocytes and macrophages in AHF.

**Female Mice Activate the Inflammation-Resolution Sensor and Cytokines in the LVI After MI**

To define cardiac healing in AHF and CHF, the differences in cytokines and the bioactive lipid mediator receptors of male and female mice and the levels of proinflammatory and inflammation controlling cytokines within the spleen and LVI at MI d1 were measured. The male mice showed increased expression of TNF-α, IL-1β, ARG-1, and MRC-1 within the spleen (Figure 8A through 8D); and the female mice showed higher expression of the same cytokines within the LVI at MI d1 (Figure 8E through 8H). The male mice showed higher protein levels of TNF-α in the LVI compared with females, indicating divergence in the levels of RNA and protein (Figure 8I and 8J). Male mice also had increased expression of HO-1 in the LVI (Figure 8K). Furthermore, we determined receptors (ALX/FPR2, EP2, and EP4) and cypoxins (EETs) metabolizing enzyme (EPHX2)
gene expression within the spleen and LVI and found minor sex-specific differences in levels of ALX/FPR2, EP4, EP2, and EPHX2 within the spleen (Figure 9A through 9D). In infarcted heart, ALX/FPR2 and EP4 activation were significant in female mice, suggestive of amplified receptor activation in female mice during cardia repair (Figure 9E through 9H).

**DISCUSSION**

After the ischemic insult, unresolved inflammation-directed HF has increasingly become one of the main causes of death, both within the United States and worldwide; however, the differences in how HF is presented and managed between men and women have been inadequately investigated. The major risk factors for CVD are universal and independent of sex and can include aging, obesity, hypertension, diabetes mellitus, alcohol consumption, and smoking.9,30 Regardless of this, women are less likely to develop CVD or HF compared with men, unless they are significantly older.30–32 In addition, hypertension has a greater impact on men developing HF than women.30 To define physiological inflammation and cardiac repair in the absence of diversified risk factors, we described leukocyte-directed molecular and cellular differences in male and female mice. Our key findings are as follows: female mice (1) have higher survival and lower rupture rates; (2) have amplified reparative monocytes, DCs, and macrophages; and (3) produced higher levels of EETs compared with male mice during the cardiac repair. Our functional and molecular analysis provides understanding about the difference between male and female mice in cardiac repair (Figure 9I).

Differences in cardiovascular morbidity and mortality caused by sex have not been adequately researched. Most mortalities develop with age, and several reports have only studied young mice in this context. Therefore, studies on phenotypical determinations need to balance sex ratios to avoid misleading information when a sex difference does exist. Our study showed the overall survival after MI is higher in female mice compared with male mice, which is in accordance with humans, where women have a
lower risk for HF than men.\textsuperscript{31,33} It was well noted that major differences in the survival rates between the male and female mice occur during AHF, whereas both male and female mice showed no difference in survival rates during CHF. This study showed female mice having a rupture rate of 7%, whereas in male mice, rupture rates were 28% in AHF. In recent decades, research on sex-related differences in cardiovascular medicine has been focused on demonstrating the role of hormones, but leukocyte phenotypes and their population differences caused by sex remained understudied.

The post-MI leukocyte-derived growth factors, cytokines, and different mediators are major contributors to inflammation and resolution. Moreover, leukocyte populations differ considerably within male and female mice during clot formation and AHF.\textsuperscript{12,34,35} Female mice showed higher population of reparative monocytes and macrophages (Ly6C\textsuperscript{lo} monocytes and Ly6C\textsuperscript{lo} macrophages), indicative of healing response.\textsuperscript{12,36} Studies have demonstrated that DC numbers increase in infarcted hearts of experimental models,\textsuperscript{37} and decreased numbers of DCs in human infarcted myocardial tissue are associated with impaired reparative fibrosis and the development of cardiac ruptures after MI.\textsuperscript{29} Female mice not only displayed high DC populations at both AHF and CHF, which provides early inflammation-resolution benefits and plays a role in fibrotic repair. DC infiltration in human infarcted myocardium showed a strong association between the number of DCs and impaired reparative fibrosis, leading to the development of cardiac rupture, and suggests a protective role of DCs during the post-MI cardiac repair process.\textsuperscript{29} In humans, women amplify D-series resolvin biosynthesis compared with men to terminate systemic inflammation induced by typhoid vaccine, which confirms sex-specific SPM differences.\textsuperscript{4} In mice, male and female equally biosynthesized SPMs with leukocyte expressed lipoxygenase activation after MI. However, although the female mice have increased inflammatory cytokines (TNF-\textgreek{a}, IL-1\textgreek{b}, ARG-1, and MRC-1) within the infarcted heart at day 1 after MI, the males have increases of the same cytokines within the spleen, indicating a difference in cardiac reaction of inflammation or reaction time between the male and female mice. This could also explain the lower rupture rates in female mice. Female mice also have a higher expression of receptors (ALX/FPR2 and EP4) within the LVI at MI-d1, indicating an increased ability to bind to both proinflammatory and proresolving ligands, supporting the

Figure 7. Female mice intensified leukocyte response compared with male mice after myocardial infarction (MI). (A) Plot showing dendritic cells (CD11c\textsuperscript{+}) at No-MI, MI-d3, MI-d5, and MI-d56 in male and female mice. (B) Histograms showing dendritic cells at No-MI, MI-d3, MI-d5 and MI-d56 in male and female mice. (C) Plot showing neutrophils (Ly6G\textsuperscript{+}) at No-MI, MI-d3, MI-d5, and MI-d56 in male and female mice. (D) Histograms showing neutrophils expression at No-MI, MI-d3, MI-d5 and MI-d56 in male and female mice. (E, F, G) Line graphs showing the percentage of the cell population of CD11c and Ly6G. (No-MI: males (n=4), females (n=4); MI-d1: males (n=5), females (n=6); \textsuperscript{1}P<0.05 vs male at respective time point; \textsuperscript{*}P<0.05 compared with no-MI naïve controls.
cardiac repair and recovery of the myocardium. Pain and inflammation are an obvious response to injury and wound healing, particularly resolvin D5 as one of the SPM moiety control chemotherapy-induced pain in females compared with males; thus, additional SPM investigation is warranted for precise and personnel sex-specific treatment.38

Cardiac healing after a heart attack is marked with overlapping inflammation and resolution processes that are initiated when leukocytes arrive at the site of infarction.12 Historically, the resolution was considered as a passive event not considered as an active program and is coordinated by immune responsive leukocyte expressed lipoxygenase activation and interaction with essential fatty acids.12,39,40 Quantitative analyses of leukocyte-directed SPMs (D-E series resolvins, protectins, maresins, and lipoxins) and cytochrome P450 epoxygenases mediated EETs suggest that male and female mice equally respond to the biosynthesis of SPMs. Of note, females biosynthesize higher levels of 11, 12 EETs compared with males. EETs manage microenvironment during inflammatory response, which favors macrophage reparative phenotype essential for cardiac healing and repair.22,41,42 The improved cardiac function and better survival in young risk-free female mice is attributable to the confounding factors associated with endogenous biosynthesis of the EET-enriched microenvironment. In this report, females mice have a longer survival with HF with a similar prevalence of HF than males. However, the outcome may change when various other risk factors (ie, processed and packed food intake, age, comedication, and metabolic defects) exist in preclinical or clinical setting.21,23,43,44 Several reports have only studied mice at a young age, and it remains unknown if sex differences occur in other studies when research extends to aging animals with overlapping risk factors, such as obesity, insulin resistance, hyperglycemia, or high blood pressure.2

**Strengths**
The presented study provided sex-specific temporal evidence with quantitative levels of bioactive lipid mediators in AHF and CHF. Survival, functional recovery, and leukocyte-directed cardiac repair that showed sex-specific signs of fine-tuned physiological inflammation.
in female mice without the risk factors that are common in a clinical setting (Figure 9I).

**Limitations and Perspective**

In women, estrogen levels and leukocyte activity during the menstrual, proliferative, and secretory phases of menstrual cycle vary and remain uncovered in mice cardiac injury experiments. In addition, the significance of some sex-related phenotype diversities warrants further investigation.

**ARTICLE INFORMATION**

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**REFERENCES**


**Figures S1–S6**

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Disclosures

None.

**Supplementary Materials**

Figures S1–S6


Supplemental Material
Figure S1. (A) Gating strategy for flow cytometry. (B-C) Donuts pie chart showing the survival rates of both sexes in acute and chronic HF.
Figure S2. Heat map showing differential quantity of specialized pro-resolving mediators (SPMs) and epoxyeicosatrienoic acid (EETs) in male and female mice.
Figure S3. Biosynthesis of arachidonic acid (AA)-derived proinflammatory mediators in male and female mice post-MI. (A) Pie chart showing the AA-derived proinflammatory mediators in the left ventricle of male mice at MI-d1. (B) Pie chart showing the AA-derived proinflammatory mediators in the left ventricle of female mice at MI-d1. (C) Bar graph showing AA-derived proinflammatory mediators of the left ventricle at MI-d1 of male and female mice. MI-d1: males (n=3), females (n=4).
Figure S4. Endogenous biosynthesis of docosahexaenoic acid (DHA) and eicosapentaenoic acid (EPA)-derived SPM precursors in male and female mice post-MI (A) Pie chart showing the DHA and EPA precursors in the left ventricle of male mice at MI-d1. (B) Pie chart showing the DHA and EPA precursors in the left ventricle of female mice at MI-d1. (C) Bar graph showing DHA and EPA precursors of the left ventricle at MI-d1 of male and female mice. MI-d1: males (n=3), females (n=4).
Figure S5. Structural changes in cardiomyocyte area. Wheat germ agglutinin (WGA) staining showing cardiomyocyte area in male and female mice. WGA = green, Hoechst = blue, [No-MI: males (n=4), females (n=4); MI-d1: males (n=5), females (n=6); MI-d56: males (n=4), females (n=4)].
Figure S6. Female mice have higher reparative macrophages (MΦ) compared to male mice post-MI. (A) Plot showing macrophages at No-MI, MI-d1, MI-d3, MI-d5, and MI-d56 in male and female mice. (B) Plot showing Ly6C<sub>lo</sub> and Ly6C<sub>hi</sub> macrophages at No-MI, MI-d1, MI-d3, MI-d5, and MI-d56 in male and female mice. (C-E) Line graphs showing the percentage of the cell population for CD11b<sup>+</sup>, Ly6C<sup>hi</sup>MΦ, and Ly6C<sup>lo</sup>MΦ. [No-MI: males (n=4), females (n=4); MI-d1: males (n=5), females (n=6); $p<0.05$ vs male at respective time point; *p<0.05 compared to no-MI naïve controls.]